ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

AZOPT 10 mg/ml eye drops, suspension

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each ml of suspension contains 10 mg brinzolamide.

Excipients:

Each ml of suspension contains 0.15 mg benzalkonium chloride.

For a full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Eye drops, suspension.

White to off-white suspension.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

AZOPT is indicated to decrease elevated intraocular pressure in:

- ocular hypertension
- open-angle glaucoma

as monotherapy in adult patients unresponsive to beta-blockers or in adult patients in whom beta-blockers are contraindicated, or as adjunctive therapy to beta-blockers or prostaglandin analogues (see also section 5.1).

4.2 Posology and method of administration

Posology

When used as monotherapy or adjunctive therapy, the dose is one drop of AZOPT in the conjunctival sac of the affected eye(s) twice daily. Some patients may have a better response with one drop three times a day.

When substituting another ophthalmic antiglaucoma agent with AZOPT, discontinue the other agent and start the following day with AZOPT.

If more than one topical ophthalmic medicinal product is being used, the medicines must be administered at least 5 minutes apart.

If a dose is missed, treatment should be continued with the next dose as planned. The dose should not exceed one drop in the affected eye(s) three times daily.

Method of administration

For ocular use.

Nasolacrimal occlusion or gently closing the eyelid after instillation is recommended. This may reduce the systemic absorption of medicinal products administered via the ocular route and result in a decrease in systemic side effects.

Instruct the patient to shake the bottle well before use. To prevent contamination of the dropper tip and suspension, care must be taken not to touch the eyelids, surrounding areas or other surfaces with the dropper tip of the bottle. Instruct patients to keep the bottle tightly closed when not in use.

Elderly population

No dose adjustment in elderly patients is necessary.

Paediatric population

The efficacy and safety of AZOPT in patients below the age of 18 have not been established and its use is not recommended in these patients. However, there is limited experience in children. The safety and efficacy of AZOPT have been studied in a small number of paediatric patients less than 6 years of age (see also section 4.4, 4.8 and 5.1).

Hepatic and renal impairment

AZOPT has not been studied in patients with hepatic impairment and is therefore not recommended in such patients.

AZOPT has not been studied in patients with severe renal impairment (creatinine clearance < 30 ml/min) or in patients with hyperchloraemic acidosis. Since brinzolamide and its main metabolite are excreted predominantly by the kidney, AZOPT is therefore contra-indicated in such patients (see also section 4.3).

4.3 Contra-indications

- Hypersensitivity to the active substance or any of the excipients.
- Known hypersensitivity to sulphonamides (see also section 4.4).
- Severe renal impairment.
- Hyperchloraemic acidosis.

4.4 Special warnings and precautions for use

Systemic effects

AZOPT is a sulphonamide inhibitor of carbonic anhydrase and, although administered topically, is absorbed systemically. The same types of adverse reactions that are attributable to sulphonamides may occur with topical administration. If signs of serious reactions or hypersensitivity occur, discontinue the use of this preparation.

Acid-base disturbances have been reported with oral carbonic anhydrase inhibitors. Brinzolamide has not been studied in pre-term infants (less than 36 weeks gestational age) or those less than 1 week of age. Patients with significant renal tubular immaturity or abnormalities should only receive brinzolamide after careful consideration of the risk benefit balance because of the possible risk of metabolic acidosis.

Oral carbonic anhydrase inhibitors may impair the ability to perform tasks requiring mental alertness and/or physical coordination in elderly patients. AZOPT is absorbed systemically and therefore this may occur with topical administration.

Concomitant therapy

There is a potential for an additive effect on the known systemic effects of carbonic anhydrase inhibition in patients receiving an oral carbonic anhydrase inhibitor and AZOPT. The concomitant administration of AZOPT and oral carbonic anhydrase inhibitors has not been studied and is not recommended (see also section 4.5).

AZOPT was primarily evaluated in concomitant administration with timolol during adjunctive glaucoma therapy. Additionally the IOP-reducing effect of AZOPT as adjunctive therapy to the prostaglandin analogue travoprost has been studied. No long term data are available on the use of AZOPT as adjunctive therapy to travoprost(see also section 5.1).

There is limited experience with AZOPT in the treatment of patients with pseudoexfoliative glaucoma or pigmentary glaucoma. Caution should be used in treating these patients and close monitoring of intraocular pressure (IOP) is recommended. AZOPT has not been studied in patients with narrow-angle glaucoma and its use is not recommended in these patients.

The possible role of brinzolamide on corneal endothelial function has not been investigated in patients with compromised corneas (particularly in patients with low endothelial cell count). Specifically, patients wearing contact lenses have not been studied and careful monitoring of these patients when using brinzolamide is recommended, since carbonic anhydrase inhibitors may affect corneal hydration and wearing contact lenses might increase the risk for the cornea. Likewise, in other cases of compromised corneas such as patients with diabetes mellitus, careful monitoring is recommended.

Benzalkonium chloride, which is commonly used as a preservative in ophthalmic products, has been reported to cause punctate keratopathy and/or toxic ulcerative keratopathy. Since AZOPT contains benzalkonium chloride, close monitoring is required with frequent or prolonged use in dry eye patients, or in conditions where the cornea is compromised.

AZOPT has not been studied in patients wearing contact lenses. AZOPT contains benzalkonium chloridewhich may cause eye irritation and is known to discolour soft contact lenses. Contact with soft contact lenses is to be avoided. Patients must be instructed to remove contact lenses prior to the application of AZOPT and wait 15 minutes after instillation of the dose before reinsertion.

Potential rebound effects following cessation of treatment with AZOPT have not been studied; the IOP-lowering effect is expected to last for 5-7 days.

4.5 Interaction with other medicinal products and other forms of interaction

Specific interaction studies with other medicinal products have not been performed with AZOPT. In clinical studies, AZOPT was used concomitantly with prostaglandin analogues and timolol ophthalmic preparations without evidence of adverse interactions. Association between AZOPT and miotics or adrenergic agonists has not been evaluated during adjunctive glaucoma therapy.

AZOPT is a carbonic anhydrase inhibitor and, although administered topically, is absorbed systemically. Acid-base disturbances have been reported with oral carbonic anhydrase inhibitors. The potential for interactions must be considered in patients receiving AZOPT.

The cytochrome P-450 isozymes responsible for metabolism of brinzolamide include CYP3A4 (main), CYP2A6, CYP2C8 and CYP2C9. It is expected that inhibitors of CYP3A4 such as ketoconazole, itraconazole, clotrimazole, ritonavir and troleandomycin will inhibit the metabolism of brinzolamide by CYP3A4. Caution is advised if CYP3A4 inhibitors are given concomitantly. However, accumulation of brinzolamide is unlikely as renal elimination is the major route. Brinzolamide is not an inhibitor of cytochrome P-450 isozymes.

4.6 Pregnancy and lactation

Pregnancy

There are no or limited amount of data from the use of brinzolamide in pregnant women. Studies in animals have shown reproductive toxicity (see also section 5.3). AZOPT is not recommended during pregnancy and in women of childbearing potential not using contraception.

Lactation

It is not known whether brinzolamide/metabolites are excreted in human milk. Animal studies have shown the excretion of brinzolamide in breast milk. Brinzolamide should only be used during breast-feeding when the benefit of breast-feeding for the child and the benefit of therapy for the woman outweigh the possible risks.

4.7 Effects on ability to drive and use machines

Temporary blurred vision or other visual disturbances, may affect the ability to drive or use machines (see also section 4.8). If blurred vision occurs at instillation, the patient must wait until the vision clears before driving or using machines.

Oral carbonic anhydrase inhibitors may impair the ability of elderly patients to perform tasks requiring mental alertness and/or physical coordination (see also section 4.4).

4.8 Undesirable effects

In clinical studies involving over 1800 patients treated with AZOPT as monotherapy or adjunctive therapy to timolol maleate 5 mg/ml, the most frequently reported treatment-related adverse reactions were: dysgeusia (5.8%) (bitter or unusual taste, see description below) and temporary blurred vision (5.8%) upon instillation, lasting from a few seconds to a few minutes (see also section 4.7).

The following adverse reactions were assessed to be treatment-related and are classified according to the following convention: very common ($\geq 1/10$), common ($\geq 1/100$) to <1/10), uncommon ($\geq 1/1,000$), rare ($\geq 1/10,000$), very rare (<1/10,000), or not known (cannot be estimated from the available data). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness. The adverse reactions were obtained from clinical trials and postmarketing spontaneous reports.

System Organ Classification	MedDRA Preferred Term	
Infections and infestations	<u>Uncommon</u> : nasopharyngitis, pharyngitis, sinusitis <u>Not Known</u> : rhinitis	
Blood and lymphatic system disorders	<u>Uncommon</u> : red blood cell count decreased, blood chloride increased	
Immune system disorders	Not Known: hypersensitivity	
Psychiatric disorders	<u>Uncommon</u> : apathy, depression, depressed mood, libido decreased, nightmare, insomnia, nervousness	
Nervous system disorders	Common: dysgeusia, headache	
	<u>Uncommon</u> : somnolence, motor dysfunction, amnesia, memory	
	impairment, dizziness, paraesthesia	
	Not Known: tremor, hypoaesthesia, ageusia	

Eva digardara	Common blomboritie blurred vision are imitation are main dry are
Eye disorders	Common: blepharitis, blurred vision, eye irritation, eye pain, dry eye,
	eye discharge, eye pruritus, foreign body sensation in eyes, ocular
	hyperaemia
	<u>Uncommon</u> : corneal erosion, keratitis, punctate keratitis, keratopathy,
	deposit eye, corneal staining, corneal epithelium defect, corneal
	epithelium disorder, intraocular pressure increased, optic nerve cup/disc
	ratio increased, corneal oedema, conjunctivitis, eye swwelling,
	meibomianitis, diplopia, glare, photophobia, photopsia, visual acuity
	reduced, allergic conjunctivitis, pterygium, scleral pigmentation,
	asthenopia, ocular discomfort, abnormal sensation in eye,
	keratoconjunctivitis sicca, hypoaesthesia eye, subconjunctival cyst,
	conjunctival hyperaemia, eyelids pruritus, eyelid margin crusting,
	eyelid oedema, lacrimation increased
	Not Known: corneal disorder, visual disturbance, eye allergy,
	madarosis, eyelid disorder, erythema of eyelid
Ear and labyrinth disorders	Uncommon: tinnitus
Eur una ladyiman alsoraets	Not Known: vertigo
Cardiac disorders	<u>Uncommon</u> : cardio-respiratory distress, angina pectoris, bradycardia,
Cardiac disorders	
	palpitations heart rate irregular
	Not Known: arrhythmia, tachycardia, hypertension, blood pressure
	increased, heart rate increased
Respiratory, thoracic and mediastinal	<u>Uncommon</u> : dyspnoea, bronchial hyperactivity, cough, epistaxis,
disorders	pharyngolaryngeal pain, throat irritation, nasal congestion, upper
	respiratory tract congestion, postnasal drip, rhinorrhoea, sneezing, nasal
	dryness
	Not Known: asthma
Gastrointestinal disorders	Common: dry mouth
	<u>Uncommon</u> : oesophagitis, diarrhoea, nausea, vomiting, dyspepsia,
	upper abdominal pain, abdominal discomfort, stomach discomfort,
	flatulence, frequent bowel movements, gastrointestinal disorder,
TT (1212 12 1	hypoaesthesia oral, paraesthesia oral
Hepatobiliary disorders	Not Known: liver function test abnormal
Skin and subcutaneous tissue disorders	<u>Uncommon</u> : urticaria, rash, rash maculo-papular, pruritus generalized,
	alopecia, skin tightness
	Not Known: dermatitis, erythema
Musculoskeletal and connective tissue	<u>Uncommon</u> : back pain, muscle spasms, myalgia
disorders	Not Known: arthralgia, pain in extremity
Renal and urinary disorders	Uncommon: renal pain
,	Not Known: pollakiuria
Reproductive system and breast disorders	Uncommon: erectile dysfunction
General disorders and administration site	<u>Uncommon</u> : pain, chest discomfort, asthenia, fatigue, feeling abnormal,
conditions	feeling jittery, irritability
Conditions	Not Known: chest pain, peripheral oedema, malaise, medication residue
Initiate poissoning and1	
Injury, poisoning and procedural	<u>Uncommon</u> : foreign body in eye
complications	

In small short-term clinical trials, approximately 12.5% of paediatric patients were observed to experience adverse reactions, the majority of which were local, non-serious ocular reactions such as conjunctival hyperaemia, eye irritation, eye discharge, and lacrimation increased (see also section 5.1).

Dysgeusia (bitter or unusual taste in the mouth following instillation) was the most frequently reported systemic adverse reaction associated with the use of AZOPT during clinical studies. It is likely caused by passage of the eye drops in the nasopharynx via the nasolacrimal canal. Nasolacrimal occlusion or gently closing the eyelid after instillation may help reduce the incidence of this effect (see also section 4.2).

AZOPT is a sulphonamide inhibitor of carbonic anhydrase with systemic absorption. Gastrointestinal, nervous system, haematological, renal and metabolic effects are generally associated with systemic carbonic anhydrase inhibitors. The same type of adverse reactions that are attributable to oral carbonic anhydrase inhibitors may occur with topical administration.

No unexpected adverse reactions have been observed with AZOPT when used as adjunctive therapy to travoprost. The adverse reactions seen with the adjunctive therapy have been observed with each active substance alone.

4.9 Overdose

No case of overdose has been reported.

Treatment should be symptomatic and supportive. Electrolyte imbalance, development of an acidotic state, and possible nervous system effects may occur. Serum electrolyte levels (particularly potassium) and blood pH levels must be monitored.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic Group: Antiglaucoma preparations and miotics, carbonic anhydrase inhibitors, ATC code: S01EC04

Carbonic anhydrase (CA) is an enzyme found in many tissues of the body, including the eye. Carbonic anhydrase catalyses the reversible reaction involving the hydration of carbon dioxide and the dehydration of carbonic acid.

Inhibition of carbonic anhydrase in the ciliary processes of the eye decreases aqueous humour secretion, presumably by slowing the formation of bicarbonate ions with subsequent reduction in sodium and fluid transport. The result is a reduction in intraocular pressure (IOP) which is a major risk factor in the pathogenesis of optic nerve damage and glaucomatous visual field loss. Brinzolamide, an inhibitor of carbonic anhydrase II (CA-II), the predominant iso-enzyme in the eye, with an *in vitro* IC $_{50}$ of 3.2 nM and a K_i of 0.13 nM against CA-II.

The IOP-reducing effect of AZOPT as adjunctive therapy to the prostaglandin analogue travoprost was studied. Following a 4 week run-in with travoprost, patients with an IOP \geq 19 mmHg were randomized to receive added treatment with brinzolamide or timolol. An additional decrease in mean diurnal IOP of 3.2 to 3.4 mmHg for the brinzolamide group and 3.2 to 4.2 mmHg for the timolol group were observed. There was an overall higher incidence of non-serious ocular adverse reactions, mainly related to signs of local irritation, in the brinzolamide/travoprost groups. The events were mild and did not affect the overall discontinuation rates in the studies (see also section 4.8).

A clinical trial was conducted with AZOPT in 32 paediatric patients less than 6 years of age, diagnosed with glaucoma or ocular hypertension. Some patients were naive to IOP therapy whilst others were on other IOP-lowering medicinal product(s). Those who had been on previous IOP medicinal product(s) were not required to discontinue their IOP medicinal product(s) until initiation of monotherapy with AZOPT.

Among patients who were naive to IOP therapy (10 patients), the efficacy of AZOPT was similar to that seen previously in adults, with mean IOP reductions from baseline ranging up to 5 mmHg. Among patients who were on topical IOP-lowering medicinal product(s) (22 patients), mean IOP increased slightly from baseline in the AZOPT group.

5.2 Pharmacokinetic properties

Following topical ocular administration, brinzolamide is absorbed into the systemic circulation. Due to its high affinity for CA-II, brinzolamide distributes extensively into the red blood cells (RBCs) and exhibits a long half-life in whole blood (mean of approximately 24 weeks). In humans, the metabolite N-desethylbrinzolamide is formed, which also binds to CA and accumulates in RBCs. This metabolite binds mainly to CA-I in the presence of brinzolamide. In plasma, both brinzolamide and N-desethylbrinzolamide concentrations are low and generally below assay quantitation limits (<7.5 ng/ml).

Binding to plasma proteins is not extensive (about 60%). Brinzolamide is eliminated primarily by renal excretion (approximately 60%). About 20% of the dose has been accounted for in urine as metabolite. Brinzolamide and N-desethylbrinzolamide are the predominant components in the urine along with trace levels (<1%) of the N-desmethoxypropyl and O-desmethyl metabolites.

In an oral pharmacokinetic study, healthy volunteers received 1 mg capsules of brinzolamide twice daily for up to 32 weeks and RBC CA activity was measured to assess the degree of systemic CA inhibition.

Brinzolamide saturation of RBC CA-II was achieved within 4 weeks (RBC concentrations of approximately 20 μ M). N-Desethylbrinzolamide accumulated in RBCs to steady state within 20-28 weeks reaching concentrations ranging from 6-30 μ M. The inhibition of total RBC CA activity at steady state was approximately 70-75%.

Subjects with moderate renal impairment (creatinine clearance of 30-60 ml/minute) were administered 1 mg of brinzolamide twice daily orally for up to 54 weeks. Brinzolamide RBC concentration ranged from about 20 to 40 μ M by week 4 of treatment. At steady-state, brinzolamide and its metabolite RBC concentrations ranged from 22.0 to 46.1 and 17.1 to 88.6 μ M, respectively.

N-desethylbrinzolamide RBC concentrations increased and total RBC CA activity decreased with decreasing creatinine clearance but brinzolamide RBC concentrations and CA-II activity remained unchanged. In subjects with the highest degree of renal impairment inhibition of total CA activity was greater although it was inferior to 90% at steady-state.

In a topical ocular study, at steady-state, brinzolamide RBC concentrations were similar to those found in the oral study, but levels of N-desethylbrinzolamide were lower. Carbonic anhydrase activity was approximately 40-70% of predose levels.

5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans with brinzolamide based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, and carcinogenic potential.

Developmental toxicity studies in rabbits with oral doses of brinzolamide of up to 6 mg/kg/day (125 times the recommended human ophthalmic dose) revealed no effect on foetal development despite significant maternal toxicity. Similar studies in rats resulted in slightly reduced ossification of skull and sternebrae of foetuses of dams receiving brinzolamide at doses of 18 mg/kg/day (375 times the recommended human ophthalmic dose), but not 6 mg/kg/day. These findings occurred at doses that caused metabolic acidosis with decreased body weight gain in dams and decreased foetal weights. Dose-related decreases in foetal weights were observed in pups of dams receiving brinzolamide orally ranging from a slight decrease (about 5-6%) at 2 mg/kg/day to nearly 14% at 18 mg/kg/day. During lactation, the no adverse effect level in the offspring was 5 mg/kg/day.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Benzalkonium chloride, mannitol (E421), carbomer 974P, tyloxapol, edetate disodium, sodium chloride, hydrochloric acid/sodium hydroxide (to adjust pH) purified water.

6.2 Incompatibilities

Not applicable.

6.3 Shelf-life

2 years.

4 weeks after first opening.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

5 and 10 ml opaque low density polyethylene bottles with polypropylene screw caps (droptainer).

The following pack sizes are available: outer cartons containing 1 x 5 ml, 3 x 5 ml and 1 x 10 ml bottles. Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Alcon Laboratories (UK) Ltd. Pentagon Park Boundary Way Hemel Hempstead Herts HP2 7UD United Kingdom.

8. MARKETING AUTHORISATION NUMBERS

EU/1/00/129/001-3

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 9 March 2000 Date of last renewal: 9 March 2005

10. DATE OF REVISION OF THE TEXT

Detailed information on this product is available on the website of the European Medicines Agency (EMEA) http://www.ema.europa.eu

ANNEX II

- A MANUFACTURING AUTHORISATION HOLDERS RESPONSIBLE FOR BATCH RELEASE
- B CONDITIONS OF THE MARKETING AUTHORISATION

A. MANUFACTURING AUTHORISATION HOLDER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

S.A. Alcon-Couvreur N.V., Rijksweg 14, B-2870 Puurs, Belgium.

or

Alcon Cusí, S.A., Camil Fabra 58, 08320 El Masnou, Barcelona, Spain.

The printed package leaflet of the medicinal product must state the name and address of the manufacturer responsible for the release of the concerned batch.

B. CONDITIONS OF THE MARKETING AUTHORISATION

CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE IMPOSED ON THE MARKETING AUTHORISATION HOLDER

Medicinal product subject to medical prescription

• CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

Not applicable

OTHER CONDITIONS

Pharmacovigilance system

The MAH must ensure that the system of pharmacovigilance, presented in Module 1.8.1. of the Marketing Authorisation, is in place and functioning before and whilst the product is on the market.

Risk Management Plan

The MAH commits to performing the studies and additional pharmacovigilance activities detailed in the Pharmacovigilance Plan, as agreed in version 01 of the Risk Management Plan (RMP) presented in Module 1.8.2. of the Marketing Authorisation and any subsequent updates of the RMP agreed by the CHMP.

As per the CHMP Guideline on Risk Management Systems for medicinal products for human use, any updated RMP should be submitted at the same time as the following Periodic Safety Update Report (PSUR).

In addition, an updated RMP should be submitted

- When new information is received that may impact on the current Safety Specification, Pharmacovigilance Plan or risk minimisation activities
- Within 60 days of an important (pharmacovigilance or risk minimisation) milestone being reached
- At the request of the EMEA

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

CARTON FOR SINGLE BOTTLE, 5 ml, 10 ml + CARTON FOR 3 x 5 ml BOTTLES

1. NAME OF THE MEDICINAL PRODUCT

AZOPT 10 mg/ml eye drops, suspension Brinzolamide

2. STATEMENT OF ACTIVE SUBSTANCE

Each ml of suspension contains 10 mg of brinzolamide

3. LIST OF EXCIPIENTS

Contains benzalkonium chloride, mannitol (E421), carbomer 974P, tyloxapol, edetate disodium, sodium chloride, hydrochloric acid/sodium hydroxide (to adjust pH) and purified water. See the package leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Eye drops, suspension;

5 ml

10 ml

 $3 \times 5ml$

5. METHOD AND ROUTE OF ADMINISTRATION

Ocular use.

Read the package leaflet before use.

Shake well before use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN

Keep out of the reach and sight of children

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8.	EXPIRY DATE		
EXD			
	EXP:		
	Discard four weeks after first opening. Opened:		
Opene			
	Opened (2): Opened (3):		
opene			
_			
9.	SPECIAL STORAGE CONDITIONS		
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR		
	WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE		
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER		
A 1			
	Laboratories (UK) Ltd.		
Pentagon Park			
	Boundary Way Hemel Hempstead		
	Herts, HP2 7UD		
	United Kingdom.		
0			
12.	MARKETING AUTHORISATION NUMBERS		
EU/1/00/129/001 1 x 5 ml			
EU/1/00/129/002 1 x 10 ml			
EU/1/0	EU/1/00/129/003 3 x 5 ml		

INSTRUCTIONS ON USE

Medicinal product subject to medical prescription.

GENERAL CLASSIFICATION FOR SUPPLY

BATCH NUMBER

13.

Lot.:

14.

15.

16 INFORMATION IN BRAILLE

azopt

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UN	ITS
BOTTLE LABEL, 5 ml & 10 ml	

1. NAME OF THE MEDICINAL PRODUCT AND ROUTE OF ADMINISTRATION
AZOPT 10 mg/ml eye drops, suspension. Brinzolamide Ocular use.
2. METHOD OF ADMINISTRATION
Read the package leaflet before use. Discard 4 weeks after first opening. Opened:
3. EXPIRY DATE
EXP:
4. BATCH NUMBER
Lot.:
5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT
5 ml 10 ml
6 OTHER

B. PACKAGE LEAFLET

PACKAGE LEAFLET: INFORMATION FOR THE USER

AZOPT 10 mg/ml eye drops, suspension

Brinzolamide

Read all of this leaflet carefully before you start using this medicine.

- Keep this leaflet. You may need to read it again.
- If you have any further questions after reading it, please ask your doctor or your pharmacist.
- This medicine has been prescribed for you. Do not pass it on to others. It may harm them even if their symptoms are the same as yours.- If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell you doctor or pharmacist.

In this leaflet

- 1. What AZOPT is and what it is used for
- 2. Before you use AZOPT
- 3. How to use AZOPT
- 4. Possible side effects
- 5. How to Store AZOPT
- 6. Further information

1. WHAT AZOPT IS AND WHAT IT IS USED FOR

AZOPT contains brinzolamide which belongs to a group of medicines called carbonic anhydrase inhibitors. It reduces pressure within the eye.

AZOPT eye drops are used to treat high pressure in the eye. This pressure can lead to an illness called glaucoma.

If the pressure in the eye is too high, it can damage your sight.

2. BEFORE YOU USE AZOPT

Do not use AZOPT

- if you have severe kidney problems.
- **if you are allergic** to any of the ingredients of AZOPT. For a full list of ingredients please see section 6
- **if you are allergic to medicines called sulphonamides.** EXAMPLES include medicines used to treat diabetes and infections and also diuretics (water tablets). AZOPT may cause the same allergy.
- if you have too much acidity in your blood (a condition called hyperchloraemic acidosis).

If you have further questions, ask your doctor for advice.

Take special care with AZOPT

Talk to your doctor

- if you have liver problems.
- if you have dry eyes or cornea problems.

- if you are taking other sulphonamide medicines

AZOPT is not to be used by people under 18 of years of age unless advised by your doctor.

Using other medicines

Please tell your doctor or pharmacist if you are taking or have recently taken any other medicines, including medicines obtained without a prescription.

If you are taking another carbonic anhydrase inhibitor (acetazolamide or dorzolamide, see section 1 WHAT AZOPT IS AND WHAT IT IS USED FOR), talk to your doctor.

Pregnancy and breast-feeding

You should not use AZOPT if you are pregnant, or might get pregnant. Talk to your doctor before you use AZOPT.

If you are breast-feeding, ask your doctor for advice.

Ask your doctor or pharmacist for advice before taking any medicine.

Driving or using machines

Do not drive or use machines until your vision is clear. You may find that your vision is blurred for a time just after using AZOPT.

AZOPT may impair the ability of elderly patients to perform tasks requiring mental alertness and/or physical coordination. If affected, take care when driving or using machines.

Important information about some of the ingredients of AZOPT

If you wear soft contact lenses. AZOPT contains benzalkonium chloride which may cause eye irritation and is known to discolour soft contact lenses. Contact with soft contact lenses is to be avoided. Remove contact lenses prior to the application of AZOPT and wait at least 15 minutes after instillation of the dose before reinsertion.

3. HOW TO USE AZOPT

Always use AZOPT exactly as your doctor has told you. You should check with your doctor or pharmacist if you are not sure.

The usual dose is 1 drop in the affected eye or eyes, twice a day-morning and night. Use this much unless your doctor told you to do something different. Only use AZOPT in both eyes if your doctor told you to. Take it for as long as your doctor told you to.

Only use AZOPT for dropping in your eyes. Turn the page for more advice.

Now turn over

3. HOW TO USE AZOPT (continued)





see side 1



2



How much to use

- Get the AZOPT bottle and a mirror
- Wash your hands
- Shake the bottle and twist off the cap
- Hold the bottle, pointing down, between your thumb and middle finger
- Tilt your head back. Pull down your eyelid with a clean finger, until there is a 'pocket' between the eyelid and your eye. The drop will go in here (picture 1)
- Bring the bottle tip close to the eye. Use the mirror if it helps
- Do not touch your eye or eyelid, surrounding areas or other surfaces with the dropper. It could infect the drops
- Gently press on the base of the bottle to release one drop of AZOPT at a time.
- **Do not squeeze the bottle:** it is designed so that a gentle press on the bottom is all that it needs (picture 2)
- After using AZOPT, press a finger to the corner of your eye, by the nose (picture 3). This helps to stop AZOPT getting into the rest of the body.
- If you take drops in both eyes, repeat the steps for your other eye.
- Put the bottle cap back on firmly immediately after use
- Use up one bottle before opening the next bottle.

If a drop misses your eye, try again.

If you get too much in your eyes, rinse it all out with warm water. Do not put in any more drops until it's time for your next regular dose.

If you forget to use AZOPT, use a single drop as soon as you remember, and then go back to your regular routine. **Do not** use a double dose to make up.

If you stop using AZOPT without speaking to your doctor, the pressure in your eye will not be controlled which could lead to loss of sight.

If you are using other eye drops, leave at least 5 minutes between putting in AZOPT and the other drops.

4. POSSIBLE SIDE EFFECTS

Like all medicines, AZOPT can cause side effects, although not everyone gets them. You can usually carry on taking the drops, unless the effects are serious.

Common side effects

(Affects 1 to 10 users in 100)

Effects in the eye: blurred vision, eye irritation, eye pain, eye discharge, itchy eye, dry eye, abnormal eye sensation, redness of the eye, eyelid itching, redness, or swelling.

General side effects: bad taste, headache, dry mouth.

Uncommon side effects

(Affects 1 to 10 users in 1,000)

Effects in the eye: increased pressure in eye, damage to the optic nerve, abnormal, double, or reduced vision, sensitivity to light, inflammation or infection of the conjunctiva, eye allergy, eye swelling, corneal disorder, inflammation of the eyelid glands, decreased eye sensation, growth on surface of eye, increased pigmentation of the eye, tired eyes, eyelid crusting, or increased tear production.

General side effects: decreased or irregular heart rate, reduced heart function, palpitations, chest pain, asthma, difficulty breathing, shortness of breath, decreased red blood cell count in blood, increased chlorine level in blood, dizziness, drowsiness, difficulty with memory, depression, difficulty sleeping, nervousness, irritability, fatigue, generalized weakness, feeling abnormal, pain, shaking, ringing in ears, decreased sex drive, male sexual difficulty, cold symptoms, chest congestion, cough, sinus infection, throat irritation, abnormal or decreased sensation in mouth, inflammation of the lining of the oesophagus, abdominal pain, nausea, vomiting, upset stomach, frequent bowel movements, diarrhoea, intestinal gas, digestive disorder, kidney pain, muscle pain, muscle spasms, back pain, nose bleeds, dry nose, runny nose, stuffy nose, sneezing, rash, abnormal skin sensation, itching, loss of hair.

Additional side effects that have been reported include:

Effects in the eye: eyelid abnormality, decreased growth or number of eyelashes.

General side effects: increased allergic symptoms, increased blood pressure, increased heart rate, abnormal liver blood tests, , frequent urination, swelling of the extremities, decreased sensation, decreased taste sensation, joint pain, pain in extremity, skin redness, inflammation, or itching.

If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell your doctor or pharmacist.

5. HOW TO STORE AZOPT

Keep out of the reach and sight of children.

Do not use AZOPT after the expiry date which is stated on the bottle and box after "EXP". The expiry date refers to the last day of the month.

This medicine does not require any special storage conditions.

You must throw away a bottle four weeks after you first opened it, to prevent infections. Write down the date you opened each bottle in the space below and in the space on the bottle label and box. For a pack containing a single bottle, write only one date.

Opened	(1):
Opened	(2):
Opened	(3):

Medicines should not be disposed of via wastewater or household waste. Ask your pharmacist how to dispose of medicines no longer required. These measures will help protect the environment.

6 FURTHER INFORMATION

What AZOPT contains

The active substance is brinzolamide 10 mg/ml.

The other ingredients are: benzalkonium chloride, carbomer 974P, edetate disodium, mannitol (E421), purified water, sodium chloride, tyloxapol. Tiny amounts of hydrochloric acid or sodium hydroxide are added to keep acidity levels (pH levels) normal.

What AZOPT looks like and the contents of the pack

AZOPT is a milky liquid (a suspension) supplied in a pack containing a 5 ml or a 10 ml plastic (droptainer) bottle with a screw cap, or in a pack containing three 5 ml plastic (droptainer) bottles with screw caps. Not all pack sizes may be marketed.

The marketing authorisation holder	Manufacturer	Manufacturer
Alcon Laboratories (UK) Ltd.,	S.A. Alcon - Couvreur N.V.,	Alcon Cusí, S.A.,
Pentagon Park	Rijksweg 14,	Camil Fabra 58,
Boundary Way,	B-2870 Puurs,	08320 El Masnou,
Hemel Hempstead,	Belgium	Spain
Herts., HP2 7UD,		
United Kingdom.		

For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder.

België/Belgique/Belgien Luxembourg/Luxemburg

SA Alcon-Couvreur NV

Алкон България ЕООД

2 + 359 2 950 15 65

2 + 32 (0)3 890 27 11 (België/Belgique/Belgien)

Nederland

Norge

Lietuva

Magyarország Alcon Hungária Gyógyszerkereskedelmi Kft.

Alcon Pharmaceuticals Ltd. atstovybė

***** + 36-1-463-9080

***** + 31 (0) 183 654321

Alcon Norge AS

Österreich

Polska

România

Slovenija

Alcon d.o.o

***** + 47 23 25 25 50

Alcon Ophthalmika GmbH

***** + 43 (0)1 596 69 70

Alcon Polska Sp. z o.o.

***** + 48 22 820 3450

***** + 370 5 2 314 756

Česká republika

България

Alcon Pharmaceuticals (Czech Republic) s.r.o.

***** + 420 225 377 333

Alcon Nederland BV

Danmark

Alcon Danmark A/S

***** + 45 3636 3434

Deutschland

Alcon Pharma GmbH

***** + 49 (0)761 1304-0

Ελλάδα

Κύπρος

Άλκον Λαμποράτορις Ελλάς ΑΕΒΕ

2 + 30 210 68 78 300 (Ελλάδα)

Eesti

Alcon Eesti

***** + 372 6 313 214

Portugal

Alcon Portugal-Produtos e Equipamentos Oftalmológicos, Lda.

***** + 351 214 400 300

2 + 386 1 422 5280

Slovenská republika

***** + 421 2 5441 0378

Alcon Pharmaceuticals Ltd – oz

S.C. Alcon Romania S.R.L **2**: +40 21 203 93 24

España

Alcon Cusí, S.A.

***** + 34 93 497 7000

France

Laboratoires Alcon

2 + 33 (0)1 47 10 47 10

Ireland

Malta

United Kingdom

Alcon Laboratories (UK) Ltd.

***** + 44 (0) 1442 34 1234 (United Kingdom)

Suomi/Finland

Alcon Finland Oy

***** +358 207 871 600

Ísland

Alcon Danmark A/S

***** + 45 3636 3434

26

Italia

 Sverige

Alcon Sverige AB ***** + 46 (0)8 634 40 00

E-post: receptionen@alconlabs.com

Latvija

Alcon Pharmaceuticals Ltd

***** + 371 7 321 121

This leaflet was last approved on

Detailed information on this product is available on the website of the European Medicines Agency (EMA) http://www.ema.europa.eu